3-Nitropropionic Acid Neurotoxicity: Visualization by Silver Staining and Implications for Use as an Animal Model of Huntington’s Disease

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The neuronal damage produced by the mitochondrial toxin 3-nitropropionic acid (3NPA) has been suggested to replicate much of the neuropathology seen in Huntington’s disease (HD) and therefore might be used in an animal model. We investigated the susceptibility to this toxin of different neuronal populations in addition to the commonly studied caudate putamen by injecting 3NPA into seven different brain regions as well as systemically. After different survival times, rats were intracardially perfused, brain sections were processed with the Gallyas silver technique, and impregnated neurons were mapped with a computerized microscope. Intracerebral administration of 3NPA resulted in a lesion, the center of which was devoid of tissue while the area was surrounded by a halo of Golgi-like impregnated neurons. In addition to local damage, rats receiving microinjections into the frontal cortex, caudate putamen, basal forebrain, and amygdala displayed argyrophilic neurons in the thalamus corresponding to the topography of thalamofugal neurons projecting to the individual injection sites. On the other hand, negligible secondary damage was seen after injections into the internal capsule, thalamus, or substantia nigra, implicating that thalamofugal axons are especially vulnerable to the local effect of this toxin. Two and a half days after systemic administration of 3NPA, a diffuse argyrophilic neuronal reaction was seen in the dorsolateral part of the caudate putamen, together with a more regionally selective staining of neurons in different cortical areas and the hippocampus. These morphopathological changes were also accompanied by motor deficits. The affected neurons in the cortical regions were primarily in those layers (V and VI) and areas (medial prefrontal, caudal insular/perirhinal, and ventral temporal) that do not project toward the lesioned striatal area; therefore, the cortical pathology may represent another primary site of action of the toxin. Among the affected neurons in the hippocampal complex were pyramidal neurons in the CA1 region as well as various neurons in the CA3 region and dentate hilar area. These studies suggest that a combination of 3NPA administration and a sensitive silver-impregnation method may unravel the potential site of primary neuronal damage in this animal model. Furthermore, these findings may contribute to the understanding of how the disease progresses in HD from the originally affected neuronal population(s) by the recruitment of closely related systems and pathways.

INTRODUCTION

Huntington’s disease (HD), an autosomal dominant disorder showing complete penetrance, results from a trinucleotide expansion on chromosome 4 (44). The common chorea seen in patients may result from the loss of enkephalin- and substance P-containing, medium-sized spiny striatal projection neurons that send efferents to the globus pallidus and substantia nigra pars reticulata (19, 32, 71). In contrast, medium-sized...
3NPA toxicity and silver staining

Aspiny interneurons containing nicotinamide adenine dinucleotide phosphate (NADPH) diaphorase, neuropeptide Y, and somatostatin and large aspiny cholinergic neurons are relatively spared (13, 19, 25, 47, 64). Furthermore, there is some evidence for damage to other brain areas, including the cortex and thalamus (20, 33, 40, 49, 53, 76).

Intrastralal or systemic administration of axon-sparing excitotoxins including kainic acid, quinolinic acid, and ibotenic acid in rodents and primates has been shown to destroy spiny striatal projection neurons and result in similar hyperkinetic movements as seen in HD (8, 17, 25, 37, 54, 74). Recently, 3-nitropropionic acid (3NPA), a plant fungal toxin which inactivates mitochondrial succinate dehydrogenase (14), has been used for studying the HD pathology. Ingestion of this toxin by eating contaminated sugar cane results in basal ganglia lesions and a marked dystonia in children (52). Similar behavioral and neuropathological changes were found in rats and mice after intrastralal and systemic administration (7, 11, 31).

Although the striatal damage produced by 3NPA in rats has been previously described (7, 12, 36, 86), the mechanism of action of this toxin is still unclear. Furthermore, whether other neuronal populations in addition to striatal neurons can be affected by 3NPA has not been examined. Therefore we injected 3NPA into areas related to basal ganglia circuits, including the frontal cortex (FC), substantia nigra, thalamus, and amygdala. We also investigated whether 3NPA could affect neurons through uptake into fibers of passage. Finally, we examined for neuronal vulnerability throughout the forebrain following systemic 3NPA administration.

The silver impregnation technique developed by Gal-lyas et al. (28) has been shown to produce a Golgi-like staining of neurons ("argyrophilic" or "dark" neurons) affected by various types of neuronal insult. These include ischemia, traumatic brain injury, and neurotoxin-induced damage (28, 29, 38), disease states which have been linked to both necrotic and apoptotic cell death (5, 24, 45, 68). This sensitive silver method was employed to detect the neuronal populations compromised by 3NPA administration as well as to follow the time course of 3NPA-induced morphopathological changes.

MATERIALS AND METHODS

Animals

Fifty adult male Sprague–Dawley rats (Zivic–Miller, Pittsburgh, PA) weighing 275–325 g were used in this study. Animals were housed on a 12/12 light/dark cycle in a temperature- and humidity-controlled room. Food and water were available ad libitum. All animal procedures were conducted in accordance with the NIH Guide for the Care and Use of Laboratory Animals.

Drug Administration

For microinjection, 3NPA (Sigma Chemical Co.) was dissolved in 0.1 M phosphate buffer. A 27-gauge needle connected via PE-20 tubing (Clay Adams) to a 5-µl Hamilton syringe was used to deliver 3NPA (500 nmol). Rats were anesthetized with sodium pentobarbital (50 mg/kg, ip) and stereotaxic (David Kopf Instruments) microinjections were aimed at the following coordinates: caudate putamen (AP +0.5, LM −2.0, DV −5.0), basal forebrain (AP −0.8, LM −2.5, DV −7.6, ipsilateral injection; or AP −0.8, LM +3.6, DV −9.7, angle = 38.5°, contralateral injection), internal capsule (AP −1.6, LM +3.0, DV −8.9, angle = 42.0°, contralateral injection), thalamus (AP −2.12, LM +3.6, DV −7.7), frontal cortex (AP +3.2, LM −0.6, DV −2.8 and AP +3.2, LM −3.8, DV −2.3), amygdala (AP −3.3, LM −4.8, DV −7.0), and substantia nigra (AP −5.3, LM −2.0, DV −7.8) according to Paxinos and Watson (65). Between four and six animals were used for each microinjection lesion group.

Injection rate was controlled by a computerized pump (Harvard Apparatus). Frontal cortex received two injections of 0.5 µl each at a rate of 0.1 µl/min and amygdala and substantia nigra received 0.4 µl at a rate of 0.1 µl/min. All other brain areas received a volume of 0.1 µl at a rate of 0.02 µl/min. Following 3NPA injections, the needle was left in place an additional 2 min before slow withdrawal from the brain. The same volume of the vehicle (0.1 M phosphate buffer) was injected into the contralateral hemisphere as controls. All rats were perfused 5 or 15 days following surgery.

For systemic injections, 3NPA injections were given at a dose of 10 mg/kg, twice daily at 12-h intervals (a total of five injections). Since many animals became severely sick or died during this treatment, the remaining animals were perfused approximately 60 h after the first injection.

To increase chances of survival and examine potential long-term effects, several animals were given a reduced dose of 3NPA ip (rather than a total of 20 mg/kg daily, only 15 mg/kg once daily was given) over a period of 3 days and were perfused 2 weeks following the end of the 3NPA treatment. Half of the animals died within the first week; the remaining animals were perfused at the end of the second week.

Perfusion and Tissue Processing

Rats were deeply anesthetized with pentobarbital and intracardially perfused with a buffered paraformaldehyde/glutaraldehyde solution (28). Brains were removed from the skull 16–24 h later, placed in the same fixative for 3 days, and then transferred to a 30% sucrose/fixative solution for an additional 3 days. Tissue sections were cut at 60 µm on a freezing microtome and were processed according to a previously described
silver technique (28). Briefly, sections were dehydrated in a graded 1-propanol series and incubated in an esterifying solution for 16 h at 56°C. Thereafter, sections were rehydrated, treated in 8% acetic acid for 10 min, and placed into a silicotungstate developer. The reaction was terminated by placing the sections in 1% acetic acid for 30 min. Adjacent sections were stained for Nissl to aid in delineating brain structures and to examine potential cell loss.

Silver-impregnated neurons were mapped using a computerized microscope and the Neurolucida software (MicroBrightField, Inc.). These Neurolucida images were then carefully aligned and overlaid on the corresponding brain maps from the Swanson atlas (79).

RESULTS

Effect of Intracerebral Microinjection of 3NPA

The locations of 3NPA microinjection sites are summarized in Fig. 1. All lesions consisted of a center of complete cell loss usually surrounded by several silver-impregnated (“dark”) neurons. In several instances we observed dark neurons outside of the vicinity of the injection, demonstrating an additional, secondary neuronal insult. Table 1 summarizes the various injection paradigms and locations of silver-impregnated neurons.

3NPA lesions in the frontal cortex (FC; Fig. 1a), caudate putamen (CP) (Fig. 1b), basal forebrain (“ipsilateral” BF lesion, Fig. 1c; “contralateral” BF injection, Fig. 1d), and amygdaloid body (Fig. 1g) resulted in secondary argyrophilic neuronal reaction in thalamic nuclei. On the other hand, injections into the internal capsule (Fig. 1e), thalamus (Fig. 1f), or substantia nigra (Fig. 1h) did not cause dark neuronal staining in areas outside the immediate vicinity of the lesion site.

The 3NPA injection into CP (Fig. 1b) was located in its medial one-third, and the microinjection needle caused minimal damage to the overlying cortex (few, if any, dark cells were observed around the needle track) and the cingulate bundle. Animals receiving striatal 3NPA microinjections displayed silver-impregnated neurons in several thalamic nuclei, including the anteroven-tral, mediodorsal, laterodorsal, and intralaminar nuclei (Fig. 2).

In the ipsilateral BF cases (Fig. 1c), the injections were centered in an area encompassing the horizontal limb of the diagonal band, substantia innominata, lateral part of the bed nucleus of the stria terminalis, and lateral hypothalamic area. The shaft of the microinjection needle penetrated the medial part of the CP, resulting in a “spillover” of the toxin and the presence of several dark neurons in this area. Affected neurons could be seen in the same thalamic nuclei as was found in striatal (CP) cases; in addition, an argyrophilic reaction was seen in the ventromedial and paraventricular thalamic nuclei. Figure 3 illustrates the distribution and characteristics of argyrophilic neurons in the thalamus in a coronal section following ipsilateral 3NPA injection into the BF. In the cases where 3NPA was injected from the contralateral hemisphere, impregnated neurons were observed in the same thalamic nuclei as after ipsilateral approach, except in the anterodorsal and ventromedial nuclei that did not contain any impregnated neurons. Also, the number of affected neurons was less after contralateral injections.

Two large 3NPA injections were directed at the same anterior–posterior level in the frontal cortex, encompassing from medial to lateral the anterior cingulate cortex, the supplemental motor area (M2), the rostral and deep parts of the lateral frontal cortex (primary motor area, M1), with a slight involvement of the prelimbic (medial prefrontal) and the dorsal agranular insular (lateral prefrontal) cortices (Fig. 1a). In addition, superficial fibers of the forceps minor (corpus callosum) and the dorsalmost part of the CP were also damaged. In these cases, silver-impregnated neurons and cellular debris were particularly noticeable in the mediodorsal, ventrolateral, and ventromedial thalamic nuclei. Additional areas containing affected cells included medial CP and globus pallidus. Figure 4 shows the distribution of dark neurons at two levels of the thalamus.

The amygdaloid injection was centered in the lateral and basolateral amygdaloid nuclei (Fig. 1g). The lesion also included the dorsal portion of the central nucleus and the caudoventral part of the putamen. Dark neu-
Many impregnated neurons showed a full Golgi-like staining of the perikaryon and dendritic processes after a short survival time (5 days); however, the presence of an uneven staining of the perikaryon (arrowhead in Fig. 3e) and corkscrew-like dendrites (asterisk in Fig. 3d) indicated an early pathological change. After a longer survival time (15 days), many neurons were in an advanced stage of degeneration, and there was neuronal debris present, characterized by shrunken and/or disintegrated cell bodies and dendrites. In other words, there were fewer fully stained neurons and more debris 15 days following CP and amygdala 3NPA injection, while in cases with FC lesions, silver-impregnated neurons and debris were observed only following this longer survival time and no staining was observed at the 5-day time point.

Since neurons or fibers passing through the injection area are also compromised resulting in anterograde degeneration of their preterminal and terminal portions, in fortuitous cases, the input to the area containing the affected neurons can also be demonstrated by applying this silver method (Fig. 5).

Animals receiving vehicle injections occasionally had only one or two dark neurons in the thalamus. There were no readily apparent motor changes observed in any of the rats receiving intracerebral injections of 3NPA and all animals survived the appropriate planned survival times.

**Effect of Systemic Injection of 3NPA**

Following two ip injections of 3NPA, rats displayed a marked dystonia and rigidity. Five systemic injections of 3NPA resulted in immobility accompanied by widespread bilateral dark neuronal staining. The most

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**TABLE 1**

Distribution of Silver-Impregnated Neurons Following Intracerebral and Systemic Injections of 3NPA

<table>
<thead>
<tr>
<th>Location of “dark” cells</th>
<th>Frontal CTX</th>
<th>Striatum</th>
<th>BFi</th>
<th>BFc</th>
<th>Amygdala</th>
<th>Systemic (ip)</th>
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<td>Laterodorsal</td>
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Note. Abbreviations: *, neuronal debris; i.s., 3NPA injection site; c, caudal; r, rostral; CTX, cortex; BFi, basal forebrain ipsilateral; BFc, basal forebrain contralateral. Rating scale: mean/unilateral structure/per section: --, no cells; +_, occasional/1 cell; +, 2–5 cells; ++, 6–10 cells; ++++, 11–20 cells; ++++++, above 20 cells.
heavily affected area was the CP, although the size of the damage was variable in each animal. Within the CP there was a spherical area occupying about two-thirds to three-fourths of the lateromedial CP and containing stained neurons, axons, and dendrites in its dense peripheral shell which was about 200 µm thick (Fig. 6a). Inside this shell only few neurons were impregnated, usually showing advanced degeneration; in addition many impregnated axons were seen especially in the bundles of the internal capsule. Nissl staining on adjacent sections revealed a reduction in cell density inside this shell of dark neurons (Fig. 6b). Parts of the lateral striatal artery inside or outside of the demarcated zone were often occluded. Although the size of this striatal injury showed some variation, the medial part of the CP and the shell of the nucleus accumbens were always free of argyrophilic neurons.

The distribution of silver-impregnated neurons following systemic 3NPA injections is shown in the series of coronal sections of Fig. 7. Particularly heavy neuronal staining can be seen in the deep layers (laminae V and VI) of the medial prefrontal cortex, the insular cortex including the claustrum, perirhinal, auditory, and ventral temporal areas. A few impregnated neurons were seen in the M1/S1 border. More caudally, particularly heavy labeling was observed in layers V and VI of all neocortical regions and the lateral entorhinal cortex. Occasionally, a few impregnated pyramidal neurons were also found in layer III (Fig. 7). Figure 8a shows the distribution of dark neurons in the parietal and temporal cortices. As can be seen, these silver-impregnated pyramidal neurons possess an abundance of spines (as shown by arrows in the figures), although the corkscrew-like dendrites suggest they are in an early phase of degeneration. Figure 9 shows further examples of silver-impregnated neurons from the temporal cortex from the same animal. Compared to neurons shown in Fig. 8, these dark neurons demonstrate a more advanced stage of degeneration, such as partial impregnation of the perikaryon (Fig. 9c) and beaded, disintegrating dendrites, and spines were no longer visible (Fig. 9b).

In cases where the dense striatal halo penetrated through the corpus callosum and insular cortex and most of the CP was involved except its most medial and caudal portions, a few impregnated neurons were also seen in the ventral part of the globus pallidus and many pyramidal neurons in layer III of the neocortical areas showed argyrophilia.

The hippocampal CA1 and CA3 fields and dentate gyrus also contained affected neurons (Fig. 10). The subiculum often contained an occluded capillary surrounded by impregnated pyramidal cell bodies. The stratum lacunosum moleculare was often densely packed with stained apical dendrites of pyramidal neurons. A few CA3c pyramidal neurons and dentate granule cells were also impregnated together with the characteristic staining of mossy fibers. Within the
FIG. 3. Distribution of affected neurons in the thalamus following injection of 3NPA into the basal forebrain from the ipsilateral side. (a) Schematic showing location of dark neurons in the thalamus. Dots represent individual silver-impregnated cells. (b) Widespread distribution of silver-impregnated cells, boxed area from (a). (c) Arrow points to a dark neuron from the boxed area in (b). (d) Dark neuron taken from (c) (arrow) under higher magnification. (e and f) Representative dark thalamic neurons. Arrowhead, uneven staining of perikaryon. Asterisk, corkscrew-like dendrite. Scale bar: b, 500 µm; c, 100 µm; d-f, 10 µm.
FIG. 4. 3NPA-induced neuronal damage in the thalamus following frontal cortex microinjection. (a and b) Diagrams of dark cells in the thalamus, plotted from corresponding coronal sections using the Neurolucida system and the Swanson atlas (79). Dots represent silver-impregnated neurons or cellular debris. (c and d) Low-magnification photographs of boxed areas in (a) and (b). (e and f) High-magnification photographs of dark cells, taken from above (c, arrow) and below (d, arrow), respectively. Asterisk, corkscrew-like dendrite. Scale bar: c and d, 100 µm; e and f, 10 µm.
patchy regions, a few inhibitory interneurons were also impregnated. Although the thalamus often contained thrombotic vessels, only a few animals showed diffuse argyrophilic reaction in the posterior portion of the lateral dorsal nucleus and adjoining posterior nuclei. This damage was similar to the dense striatal spherical staining of cells and dendrites and axons within the halo.

To examine the long-term effects of systemic 3NPA administration, several animals were given the reduced dosage of 3NPA ip (to increase chances of survival) over a period of 3 days and were perfused 2 weeks following the end of the 3NPA treatment. On Day 4, all animals showed visible motor deficits and were given sc glucose injections to improve chances of survival. Half of the animals died within the first week, but by the end of the second week, the remaining half appeared to possess normal motor abilities. Following perfusion and dark neuronal staining, there were only a few cells in the CP, cortex, and hippocampus, a drastic reduction in staining seen compared with the animals receiving the higher dose of 3NPA.

**DISCUSSION**

We have examined the neurotoxic damage induced by intracerebral and intraperitoneal delivery of 3NPA in rats, screening for neuronal populations that are vulnerable to this toxin. Our studies provide evidence that (1) following intracerebral administration of 3NPA, thalamofugal neurons are especially sensitive to this toxin when delivered into their potential target areas; (2) following systemic administration of 3NPA, in addition to the known striatal damage, a substantial number of cortical and hippocampal neurons are also compromised; and (3) the Gallyas silver-impregnation technique proved to be a sensitive method for defining the precise location and the time course of morphopathological changes of affected neurons.

**Neurotoxic Risk Assessment**

Different animal models of HD are useful for revealing the potential primary site of neuronal damage as well as for understanding the disease progression. The superior sensitivity of silver techniques to detect morphopathological changes in the brain compared to traditional Nissl staining has been known for decades (9). An example of the dramatic difference between 3NPA-induced striatal cell damage as revealed by the Gallyas silver method and Nissl staining can be observed by comparing Fig. 6a with Fig. 6b. Furthermore, smaller differences are still readily apparent after silver staining, such as with the silver-impregnated cortical neurons affected by 3NPA (Figs. 7–9); these impaired neurons in a similar experimental paradigm using Nissl staining could not be demonstrated (7). Although silver staining was recently used in combination with 3NPA administration (4), due to the fact that the particular silver technique was developed primarily for detecting degeneration of axon terminals (27), the consequent lack of staining of the somatodendritic domain prevented firm conclusions to be drawn as to identity of neuronal populations affected by 3NPA.
FIG. 6. Striatal neuronal damage following intraperitoneal administration of 3NPA. (a) Dark neuron-stained section showing dense, bilateral circles of silver-impregnated neurons and fibers in the caudate putamen. (b) Nissl-stained section showing bilateral circular areas of reduced staining (area inside dotted lines) in the caudate putamen.
Pathological conditions such as head trauma, ischemia, insulin hypoglycemia, status epilepticus, and various neurotoxin-related damage seem to induce the same type of morphopathological changes that can be detected by a recently developed silver method of Gallyas (9, 28, 29, 38, 81). The similarity of the argyrophilic reaction of neurons of various origin after different types of injury suggests a common final cellular mechanism, the nature of which remains to be established. The reliable and predictable induction of somatic argyrophilia in specific neurons, the ability to visualize a large part of the dendritic tree that allows the classification of neuronal subtypes, the simultaneous staining of the projection axons of affected neurons, and the ability to follow the progression of pathological changes of affected neurons make this new Gallyas silver technique especially attractive in the evaluation of experimental models of neurodegenerative diseases.

Evaluation of Morphopathological Changes after Intracerebral 3NPA Injection

All 3NPA injection sites displayed a complete loss of tissue in the immediate vicinity of the injection and were surrounded by several dark cells. In addition, pronounced dark neuronal staining was observed in several thalamic nuclei, remote from the injection sites. There was negligible, if any, cell staining observed in control cases.

In the cases where 3NPA was injected into the FC, compromised neurons were found in the mediadorsal,
FIG. 8. Cortical neuronal damage following intraperitoneal injection of 3NPA. (a) Diagram showing distribution of labeled cells in the cortex at the level shown in Fig. 7f. Dots represent individual silver-impregnated cells. (b) Dark neurons in perirhinal cortex. (c) Dark neurons in the parietal cortex. (d) Dark pyramidal neuron from perirhinal cortex. (e) Dark pyramidal neuron from parietal cortex located in the boxed area in (c). Arrows, dendritic spines. Scale bar: b and c, 100 µm; d and e, 10 µm.
FIG. 9. Characteristics of silver-impregnated neurons in the temporal cortex following systemic administration of 3NPA. (a) Schematic approximately 5.6 mm posterior from the bregma. (b) Dark neurons taken from boxed area in (a). (c) Dark neuron at higher magnification, taken from (b) (filled arrow). Note the uneven dark neuronal staining of the perikaryon. (d) Dark neuron at higher magnification, taken from (b) (arrowhead). Note the full silver impregnation of perikaryon and processes. (e) Dark neuron at higher magnification, taken from (b) (open arrow). Note the beaded, disintegrating dendrites. Scale bar: b, 100 µm; c-e, 10 µm.
ventromedial, and ventrolateral thalamic nuclei, which are known to project to the anterior cingulate, to the medial precentral areas, and especially to the frontal motor cortex, areas that were all damaged by the two large injections into the frontal cortex (1, 15, 22, 34, 41, 46, 48, 70). However, we did not find evidence of compromised cells in the intralaminar nuclei, thalamic areas which project sparsely to the FC (10, 35, 46). Striatal 3NPA injection (Fig. 1b) resulted in dark neuronal staining in the intralaminar thalamic nuclei, which are known to project heavily to the striatum (10, 60, 67, 78, 83). Injection of 3NPA into the amygdala showed another distinct pattern of remote silver impregnation: dark neurons were observed among thalamic areas known to project to the amygdaloid body and posterior part of the CP (50, 63, 80). Also, the strong labeling in the paraventricular thalamic nucleus after ipsilateral BF injection is compatible with the heavy termination of paraventricular axons in the bed nucleus of stria terminalis, medial striatal areas adjacent to the lateral ventricle, and the nucleus accumbens (34, 58), regions which were heavily involved at the injection site. The above cases strongly suggest that thalamofugal neurons can be specifically affected by application of 3NPA into their terminal areas.

Whether this toxin can also affect neurons through damaged axons of passage is unclear; however, a number of observations suggest such a possibility. Fibers from the anterior thalamic nuclei pass through the medial part of CP and the cingulate bundle to terminate in different portions of the cingulate cortex (21, 75). The ipsilateral BF case resulted in a spillover of the 3NPA into the medial portion of the CP and the shaft passed through the cingulate bundle, which may account for the heavy neuronal staining in the anterior thalamic nuclei compared to the contralateral BF cases. Also, the labeling in the anterior nuclei in striatal injections may be due to 3NPA uptake into fibers of passage. In a similar fashion, fibers from the reuniens thalamic nucleus pass through the rostral part of the CP and medial compartment of the cingulate bundle as they travel toward the medial cortical surface (85). Dark neurons found in the laterodorsal thalamic nucleus in the striatal and ipsilateral BF cases may relate to damage of their axons as they pass through the CP and the deep white matter to enter into the cingulate cortex (82). In some cases, however, it is difficult to determine whether the observed labeling is due to the effect of the toxin at the terminal zone or through the preterminal portion of the axon. For example, it is well known that the mediodorsal thalamic nucleus sends projections to the prefrontal cortex and these fibers are passing through the medial portion of CP (34); however, there is not a general consensus in the literature (34, 69) whether mediodorsal fibers also terminate in the CP. On the other hand, the lack of labeling of the anterior, intralaminar, and rhomboid thalamic nuclei in FC cases, or the quantitative difference in the labeling of paraventricular thalamic cells between ipsilateral and contralateral BF cases, suggests that the labeling of the parent cell bodies may depend on a dense arborization of their terminals and/or injury of their axons in the

![FIG. 10. Silver-impregnated cell bodies and dendritic processes in the hippocampus. (a) Low-magnification photomicrograph showing extensive involvement of the CA1 area. (b) Higher magnification of the boxed area of (a). Note the dense silver staining of the apical dendrites of pyramidal neurons in the stratum lacunosum-moleculare. (c) High-magnification view of apical dendrites of pyramidal cells from box in (b). Scale bar: a, 1 mm; b, 250 µm; c, 50 µm.](image)
injection area. Perhaps a combined injection of 3NPA and a retrograde tracer, such as rhodamine-labeled latex beads that are known to be taken up less often by fibers of passage, needs to be done in order to determine the potential damage to fibers of passage by this toxin. Since there was no dark neuronal staining of cortical neurons after 3NPA injection into the internal capsule (Fig. 1e), heavily myelinated fibers may be less susceptible to this toxin.

Evaluation of Pathological Changes after Systemic 3NPA Administration

In addition to dark neuronal staining in the CP following ip injection, we also observed silver-impregnated cells in deep layers of various allo- and neocortical areas, including the hippocampal formation. The involvement of the deep cerebral cortex, especially in its lateral aspect subjacent to the rhinal sulcus, was previously noted after short survival of 3NPA treatments in rats, although the nature of this pathology was not described in detail perhaps due to the use of conventional hematoxylin and eosin (HE) staining (36). The medial prefrontal cortex, where many silver-impregnated neurons were detected, projects mainly to the medial part of the CP, including the nucleus accumbens and basal forebrain areas rich in corticopetal projection neurons (30, 43, 73, 88), territories which were free of diffuse neuronal insult. The insular cortex, another region which contained high number of argyrophilic neurons, projects primarily to the fundus striati, bed nucleus of the stria terminalis, amygdala, ventro-posterior parvocellular thalamic nucleus, and brainstem visceromotor and viscero sensory nuclei (87). Finally, the perirhinal/ventral temporal cortex is considered a multimodal association area which projects to the amygdala (80). On the other hand, relatively few affected neurons were located in sensorimotor cortical areas, regions which project heavily to the lateral part of the CP (23, 55). In summary, the cortical neurons mostly affected in the systemic 3NPA cases with smaller striatal lesion do not seem to be part of the sensorimotor cortico striatal circuits but rather may be connected to limbic and autonomic centers and thus could represent an independent site of action of the toxin. On the other hand, in cases where a large proportion of the CP was included into the lesioned area, the impregnation of layer III pyramidal neurons may be the consequence of the primary striatal injury.

Potential Recovery and Selective Vulnerability of Neurons Following 3NPA Treatment

Unquestionably, the somatodendritic debris at the intracerebral injection sites as well as the shrunken, disintegrated neurons in the CP and some cortical areas after intraperitoneal 3NPA administration represents cell death as revealed by the loss of Nissl staining in adjacent sections. Whether or not the impregnated neurons at remote places from the intracerebral injections will eventually die or are simply transiently affected is a question of great importance, especially in evaluating the consequences of neuronal insult. Also, animals given the lower dose of 3NPA ip and longer survival period showed negligible dark neuronal staining and subsequent motor deficits, and, therefore, some of the staining may reflect a transient compromise of neurons, confirming a similar suggestion given by Allen et al. (4).

It is unclear why there are such localized lesions in the brain following ip 3NPA injection even though mitochondrial succinate dehydrogenase activity is reduced uniformly in all brain regions (31). Selective vulnerability of neurons may relate to biochemical composition, lack of specific neurotrophic support, substantial glutamatergic input, altered excitatory amino acid receptor subtypes, and neural system-specific expression of genetic defects (2, 51, 56). Thalamic neurons are especially vulnerable after intracerebral administration of 3NPA, since corticofugal, striatonigral, and nigrostriatal neurons were not affected when 3NPA was injected into the internal capsule, the substantia nigra, or the CP, respectively, confirming other studies (4, 66). Moreover, we could not find dark neurons in the thalami after similar large injections of kainic acid or N-methyl-o-aspartate (NMDA) into the frontal cortex (unpublished observations) which would suggest that 3NPA affects neurons both directly at the level of the cell body and at the axonal domain. This is compatible with the observation that thalamic neurons show retrograde degeneration due to damage of both their afferents and their parent axons (16).

The mechanism of striatal cell death after systemic administration of 3NPA is not well understood, although several factors may contribute to selective vulnerability of medium-sized spiny striatal neurons. Decortication, which removes putative glutamatergic inputs to the CP, can protect against 3NPA toxicity in the CP (6, 26), and NMDA receptors are upregulated in both cortex and CP following 3NPA injection (86). These are both lines of evidence for a secondary excitotoxic mechanism, in which endogenous glutamate can become toxic following a compromise in the cell’s energy levels (2, 62). However, NMDA antagonists such as MK-801 do not prevent 3NPA-induced neuronal damage (6), which suggests that 3NPA can produce neurodegeneration in both an NMDA receptor-dependent and independent manner. The sparing of striatal nitric oxide synthase (NOS)-containing cells and the protection by NOS inhibitors against striatal lesions produced by systemic administration of 3NPA suggest a role of nitric oxide in 3NPA-induced neuronal degeneration (72). Since the tricarboxylic acid cycle of GABAer-
gic neurons is selectively inhibited by 3NPA (39), it is reasonable that the degeneration is primarily localized in the CP where the majority of neurons use GABA as their transmitter. Finally, a special vulnerability is given by the lateral striatal artery, by which, under conditions such as intoxication and energy deficiency, serum proteins from thrombosis and perivascular leakage may contribute to neuronal death that localizes specifically in the lateral CP (61). The hippocampal CA1 and CA3c pyramidal neurons that we have found affected by ip 3NPA seem to be the same neurons that are highly vulnerable to hypoxic insult (42). Future studies aimed at identifying the specific neurochemical composition and connectivity of degenerating neurons in 3NPA toxicity could reveal a more precise mechanism of the selective vulnerability of these neuronal populations.

3NPA and Modeling HD

Previously, other toxins have been used in modeling HD, including kainic acid, ibotenic acid, and quisqualic acid (8, 17, 25, 37, 54, 74). 3NPA has been introduced most recently in modeling HD, due to its destructive nature in both humans and animals which results in behavioral and neuropathological features comparable to those seen in HD (7, 52, 86). Additionally, energy metabolism may be compromised in HD patients (6), and since 3NPA is a mitochondrial toxin, perhaps a similar environment in which cellular energy levels are compromised can be duplicated. NOS/NADPH–diaphorase cells are selectively spared in HD (7) as well as after 3NPA treatment (11, 72). A final attractive characteristic of 3NPA that supports its use in modeling HD is its similar age-dependent nature; children are more susceptible to its toxic effects (52), similar to the clinical symptomatology which is more pronounced in patients with young-age onset than in those with the late-onset variant of the disease (59).

As a potential model of HD, our data are in agreement with several other studies suggesting that systemically administered 3NPA may be a better model of HD than intracerebral microinjection, given the simultaneous presence of striatal pathologies and motor deficits similar to what exists in HD (24). However, systemic administration induces marked variations in the neuropathological involvement of the CP in different species. The rodent systemic model showing neuropathology in the dorsolateral CP mimics well some of the motor deficits of HD. On the other hand, a recently developed chronic systemic 3NPA model in primates with a heavy involvement of the caudate nucleus (64) replicates more the cognitive deficits of HD. The expression of different symptoms in these animal models are in line with the differential roles of the multiple prefrontostriatal circuits in motor and cognitive functions (3). Since the early destruction of the periventricular por-
thalamus is among the structures affected in HD and whether this is a consequence of the cortical or striatal pathology or represents an independent process. Similarly, given the differential vulnerability of the various hippocampal sectors to hypoxic damage (42) and the speculations of bioenergetic defects and excitotoxic-induced neurodegeneration in HD (2, 6), together with our current findings, there is the possibility of hippocampal insult in HD patients. It is hoped that further experimental studies with 3NPA using a combination of behavioral testing and a careful monitoring of the time course of degeneration in various neuronal systems may dissect the complex chain reaction leading to motor and cognitive symptoms seen in HD.

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